Generate Collection

L3: Entry 4 of 9

File: JPAB

Aug 4, 1988

PUB-NO: JP363188623A

DOCUMENT-IDENTIFIER: JP 63188623 A

TITLE: UBIDECARENONE PREPARATION HAVING IMPROVED ABSORPTION

PUBN-DATE: August 4, 1988

INVENTOR-INFORMATION:

NAME

COUNTRY

OZAWA, YASUO YAMADA, KENJI AKIMOTO, MASAYUKI TANAKA, YOSHITAKA

ASSIGNEE-INFORMATION:

NAME

COUNTRY

TAISHO PHARMACEUT CO LTD

APPL-NO: JP62021378

APPL-DATE: January 31, 1987

INT-CL (IPC): A61K 31/12; A61K 31/12; A61K 47/00; C07C 50/28

ABSTRACT:

PURPOSE: To obtain a preparation for oral administration, by adding a middle- chain fatty acid monoglycerin ester to ubidecarenone.

CONSTITUTION: 1pt.wt. ubidecarenone is blended with 0.5∼150pts.wt. middle-chain fatty acid monoglycerin ester (e.g. capric acid, caproic acid, caprylic acid, etc.) to give a preparation for oral administration having high absorption of ubidecarenone. The middle-chain fatty acid monoglycerin ester may be mixed with a third component such as vegetable oil. etc., and used as a mixture. The blending ratio of the third component is preferably 0.2∼1pt.wt. based on lpt.wt. of the ester.

COPYRIGHT: (C) 1988, JPO&Japio

Generate Collection

L6: Entry 12 of 13

File: DWPI

Aug 4, 1988

DERWENT-ACC-NO: 1988-260484

DERWENT-WEEK: 198837

COPYRIGHT 2003 DERWENT INFORMATION LTD

TITLE: Absorption-improved ubidecarenone prepn. - contg. ubidecarenone and middle chain fatty acid mono:glycerine ester(s), pref. e.g. capric acid

PATENT-ASSIGNEE:

ASSIGNEE

CODE

TAISHO PHARM CO LTD

TAIS

PRIORITY-DATA: 1987JP-0021378 (January 31, 1987)

PATENT-FAMILY:

PUB-NO

PUB-DATE

LANGUAGE

PAGES MAIN-IPC

JP 63188623 A August 4, 1988

003

APPLICATION-DATA:

PUB-NO

APPL-DATE

APPL-NO

DESCRIPTOR

JP 63188623A January 31, 1987

1987JP-0021378

INT-CL (IPC): A61K 31/12; A61K 47/70; C07C 50/28

ABSTRACTED-PUB-NO: JP 63188623A

BASIC-ABSTRACT:

An oral prepn. contg. ubidecarenone and middle-chain fatty acid monoglycerine esters.

Specifically pref. proportion of middle-chain fatty acid monoglycerin ester and ubidecarenone is 0.5-150 pts. wt. and 1 pts. wt. respectively. Pref. middle-chain fatty acid is capric acid, caproic acid, caprylic acid, etc. Plant oils can be added as a third component. Pref. their proportion to middle-chain fatty acid monoglycerin ester (1 pts. wt.) is 0.2-1 pts. wt.

USE/ADVANTAGE - Ubidecarenone, also called Coenzyme Q10, is used for cardiac incompetence and improvement of cardiac functions. However, ubidecarenone is hardly sol. in water and absorption when administered orally is bad. In order to improve absorption, ubidecarenone soft capsule is developed, but its absorptivity is still insufficient. This invention presents a new ubidecarenone oral prepn. whose absorptivity is good.

CHOSEN-DRAWING: Dwg.0/0

TITLE-TERMS: ABSORB IMPROVE UBIDECARENONE PREPARATION CONTAIN UBIDECARENONE MIDDLE CHAIN FATTY ACID MONO GLYCEROL ESTER PREFER CAPRIC ACID

ADDL-INDEXING-TERMS: COENZYME

DERWENT-CLASS: B05

CPI-CODES: B10-A06; B10-E04C; B12-F01B; B12-M11;

CHEMICAL-CODES:

Chemical Indexing M2 *02*
 Fragmentation Code
 H4 H402 H482 H8 J0 J011 J2 J271 M210 M216
 M220 M221 M222 M223 M224 M225 M231 M262 M281 M313
 M321 M332 M343 M383 M391 M416 M431 M620 M782 M903
 M904
 Markush Compounds
 198837-14901-M
 Registry Numbers
 3102R 1678D

Chemical Indexing M6 *03* Fragmentation Code M903 P522 R031 R111 R280 R301 Registry Numbers 3102R 1678D

SECONDARY-ACC-NO: CPI Secondary Accession Numbers: C1988-116001

Generate Collection

L6: Entry 8 of 13

File: JPAB

Aug 4, 1988

PUB-NO: JP363188623A

DOCUMENT-IDENTIFIER: JP 63188623 A

TITLE: UBIDECARENONE PREPARATION HAVING IMPROVED ABSORPTION

PUBN-DATE: August 4, 1988

INVENTOR - INFORMATION:

NAME

COUNTRY

OZAWA, YASUO YAMADA, KENJI AKIMOTO, MASAYUKI TANAKA, YOSHITAKA

ASSIGNEE-INFORMATION:

NAME

COUNTRY

TAISHO PHARMACEUT CO LTD

APPL-NO: JP62021378

APPL-DATE: January 31, 1987

INT-CL (IPC): A61K 31/12; A61K 31/12; A61K 47/00; C07C 50/28

ABSTRACT:

PURPOSE: To obtain a preparation for oral administration, by adding a middle- chain fatty acid monoglycerin ester to ubidecarenone.

CONSTITUTION: lpt.wt. ubidecarenone is blended with 0.5∼150pts.wt. middle-chain fatty acid monoglycerin ester (e.g. capric acid, caproic acid, caprylic acid, etc.) to give a preparation for oral administration having high absorption of ubidecarenone. The middle-chain fatty acid monoglycerin ester may be mixed with a third component such as vegetable oil. etc., and used as a mixture. The blending ratio of the third component is preferably 0.2∼1pt.wt. based on lpt.wt. of the ester.

COPYRIGHT: (C) 1988, JPO&Japio

卵日本国特許庁(JP)

⑩特許出願公開

四公開特許公報(A)

昭63-188623

識別記号 ABN ACY 庁内整理番号 7330-4C

❸公開 昭和63年(1988)8月4日

C 07 C 50/28

ACX 314

E-6742-4C

審査請求・未請求 発明の数 1 (全3頁)

❷発明の名称 吸収改善ユビデカレノン製剤

到特 顧 昭62-21378

❷出 願 昭62(1987) 1月31日

@発 明 者 小 沢 康 雄 **砂鞋** 明者 Ш BB 震 귤 **砂**発明者 秋 元 雅之 **伊** 明 者 田中 夢 孝 砂出 翔 人 大正製薬株式会社 20代 理 人 弁理士 北川 宮澄

東京都登島区高田3丁目24番1号 大正製業株式会社內東京都登島区高田3丁目24番1号 大正製業株式会社內東京都登島区高田3丁目24番1号 大正製業株式会社內東京都登島区高田3丁目24番1号 大正製業株式会社內東京都登島区高田3丁目24番1号

東京都曼島区高田3丁目24番1号

明 细·音

1. 益明の名称

妖权改善ユピデカレンン製剤

2.特許請求の範囲

ユピデカレノンに中鉄脂肪酸モノグリセリンエステル類を採加すること特徴とする経口投与製剤。

3. 是明の詳細な以明

[産業上の料用分野]

本発明は経口吸収性に優れたユピデカレノン異 剤に関する。

[健康の技術]

ユピデカレノンはコエンザイムQioまたはユピキノン10とも呼ばれるキノリン諸事体であり、 ひ不念思者の処理動態の改善、心能能低下の予防 および鬱血性心不全に併う呼吸困難、浮顔などの 改善に有効な医薬として広く使用されている。しかし、この化合物は水に極めて溶け無く、従ってこれを経口投与したときの吸収性に難点があった。そのため、特に固形製剤などでは内限した場合、消化液中への分散が悪く、吸収性に悪影響を及ぼしていた。

こうした問題点を解決する目的でユビデカレノン を抽動類に溶解または分散させた数カプセル剤が開発、溶解されている。

[発明が解決しようとする問題点]

しかしながら、この様な市販の飲力プセル剤も 吸収性において機足し得るものではなく、さらに 吸収性に扱れた銀剤の開発が望まれている。

[周囲点を解決するための手段]

本発明者らは前記問題点に増み、ユビデカレノン合有観期の経口吸収性を高めるべく、 録意検討 した結果、ユビデカレノンに中額脂肪酸モノブリ セリンエステル類を緩加した経口役与製剤が上記 目的を達成することを見出し、本発明を完成し

本発明において、中級関節酸モノグリセリンエ ステル類のユビデカレノンに対する低加部合は特 に限定されないが、通常ユピデカレノン1重量部 に対して中鎮間防酸モノグリセリンエステル質 0.5~150重量部である。

中級殷助職モノグリセリンエステル類の中鉄服 助機として好ましいものとしてはカブリン酸、カ ブロン散およびカプリル酸などである。

また、上記中質用助像モノグリセリンエステル 類に彼り前などの第三成分を抵加して混合物とし ても良い。この討合も特に限定されないが、政常 中類皮肪酸モノグリセリンエステル乗1重量部に 対して、抗三成分は0.2~1度量部であること が好ましい。

[発明の効果]

本発明によりユビデカレノンの鉄収性を高めた 盛口投与量剤を提供することができる。

字路梯5

ユピデカレノン10歳量部とカプリン酸モノグ りセリンエステル10重量部を超和、加温市解 し、この疳液をヒドロキシブロビルセルロース2 0 武造隊、結集セルロース30並量部および乳糖 29度量節の混合物に均一に分放させた。

次いで、この分数物を乾燥型粒後、ステアリン 酸マグネシウム1重量部を混合し、1錠100m の殺剤を圧縮成形した。

以験例1

(战勋勤物)

、試験実施質日より絶食させたピーグル犬(体理 10~18kg) + 1 7 3 項用いた。

(检体)

以下のカプセル剤を検体とした。

費したもの(1カプセル中ユビデカレノ ン10年合有)。

校体2:市景飲カプセル前(1カプセル中ユビ デカレノン10電合有。基剤としてプロピレング

【安施例】

以下、突返得および試験側を挙げて本発明を具 体的に延明する。

软饰细 1

. ユピデカレノン1gをカプリン酸モノグリセリ ンエステル149gに加益溶解して、ユビデカレ ノン0.67%治液を調製した。

ユビデカレノン1gもカプリン酸モノグリセリ ンエステル:大豆油=1:1の混合液149gに 加亜溶解して、ユビデカレノン0.67%溶液を

安施例3

ユビデカレノン1重量部モカブリン酸モノグリ セリンエステルタ遺量部に加進熔解し、常法によ り飲カプセル剤を問盤した。

安施例 4

カプリン酸モノグリセリンエステル30点量部 にユビデカレノン1度量節を加え、加湿熔解し て、常法により0号カブセルに充塡した。

リコールジカブリン酸を使用していた。) ・ (投与方法)

ピーグル大に各枚体(ユピデカレノン20吨/ 匹)を経口投与し直接に水30㎡を強制的に投与

(試験方法)

Lt.

血液試料の採取と処理

校体投与直前、投与後1時間、2時間、3時 間、4時間、5時間、7時間および24時間ごと に前腕が脈より血液 5 嘘を採取し、達心分離後の 血漿を放料とした。

定量法

各血型中のユピデカレノン渡皮は高速液体クロ マトグラフィー依により樹定した。[小択ら、ア ルフナイミツテル フォルシュング(Arsocia-検体1:実施例1の組成物を従カプセル形に充 Forsch)第38巻、第689頁、1986年] すなはち、血漿 0 . 5 ml に 蒸留水 0 . 5 ml を加 え、エクノール・ヘキサン混紋(2:5)7㎡で 幼山した。次にヘキサン暦 4 皿を意発乾因し、森 造に粉硫酸 0 . 5 或と 2 %塩化第二數 0 . 5 或を

(以致助果)

> 本発明の観測は対照技体よりも優れた疑口吸収 を示した。

要1 各検体投与後の血質

執体	ユビデカレノンの血吸中濃度:単位/個/四									
	1時間後	2時間後	3時間後	4時間後	5時間後	7時間接	2 4 時間接			
1	0.04	0.33	0.50	0.43	Q.44	0.44	0.22			
	(±0.01)	(±0.02)	(±0.02)	(±0.07)	(±0.09)	(±0.08)	(±0.69)			
2	0.05	0.15	0.15	0.13	0.07	0.15	0.05			
	(±0.01)	(±0.03)	(±0.03)	(±0.04)	(±0.02)	(±0.06)	(±0.02)			

()内比概準偏差

PTO: 2003-2536

Japanese Published Unexamined Patent Application (A) No. 63-188623, published August 4, 1988; Application Filing No. 62-21378, filed January 31, 1987; Inventor(s): Yasuo Ozawa et al.; Assignee: 62-21378; Japanese Title: Absorption-Improved Ubidecarenone Tablets

ABSORPTION-IMPROVED UBIDECARENONE TABLETS

CLAIM(S)

Tablets for oral intake characterized in that a middle fatty acid monoglycerin ester group is added to ubidecarenone.

DETAILED DESCRIPTION OF THE INVENTION

(Field of Industrial Application)

The present invention pertains to ubidecarenone tablets.

(Prior Art)

Ubidecarenone is a quinoline derivative generally called coenzyme Q10 or ubiquinone 10, and is widely used to improve blood circulation of patients with heart disease and to improve a respiratory problem caused by a defective heart function. This compound is not easily dissolved in water, so its absorption was a problem when orally taken in. Therefore, when it is processed into solid tablets and orally taken in, the dispersion into a digesting fluid is poor and is not absorbed well.

To solve such a problem, there has been developed and marketed soft gel capsules wherein ubidecarenone is dissolved or dispersed in a fatty group.

l

(Problems of the Prior Art to Be Addressed)

Even with these market-sold soft gel capsules, they were not totally satisfactory in absorption, and it has been demanded to develop tablets having more excellent absorption.

(Means to Solve the Problems)

The examiners of the present invention, taking the aforementioned problems into consideration, studied assiduously how to improve the absorption of ubidecarenone-containing tablets to be orally taken in. As a result, orally taken in tablets containing a middle chain fatty acid monoglycerine ester gourp in ubidecarenone can satisfy the aforementioned purpose and produced the present invention.

The ratio of the added middle chain fatty acid monoglycerine ester group relative to ubidecarenone is not specifically limited, but generally, to 1 part/weight of ubidecarenone, the middle fatty acid monoglycerine ester group is added by 0.5 – 150 parts/weight.

The preferred middle chain fatty acids out of a middle chain fatty acid monoglycerine ester group are caprylic acid, capric acid, and caproic acid.

It is also possible that a third component such as a vegetable oil is added to said middle chain fatty acid monoglycerine ester group to make an admixture. The mixing ratio in this case needs not be specified, but generally, the third component

is preferably 0.2 -1 part/weight relative to 1 part/weight of middle fatty acid monoglycerine ester group.

(Advantage)

The present invention can present tablets for oral intake that have improved absorption of ubidecarenone.

(Embodiment)

The present invention is explained below with reference to the embodiment example and test sample.

(Embodiment Example 1)

1 g of ubidecarenone was heated and dissolved in 149 g of monoglycerin caprate and a 0.67% ubidecarenone solution was thus prepared.

(Embodiment Example 2)

1 g of ubedecarenone was heated and dissolved in 149 g of admixture of monoglycerin caprate ester and soy beans oil with the mixing ratio 1:1 to prepare the ubidecarenone 0.67% solution.

(Embodiment Example 3)

1 part/weight of ubidecarenone was heated and dissolved in 9 parts/weight of monoglycerin caprate ester to prepared soft gel capsules by a conventional method.

(Embodiment Example 4)

1 part/weight of ubidecarenone was added to 30 parts/weight of monoglycerin caprate ester and heated and dissolved. This admixture was filled in No. 0 soft gel capsules by a conventional method.

(Embodiment Example 5)

10 parts/weigh of ubidecarenone and 10 parts/weight of monoglycerin caprate ester were mixed and dissolved by heat. This admixture solution was evenly dispersed in an admixture of hydroxypropyl cellulose 20 parts/weight, crystalline cellulose 30 parts/weight, and of lactose 29 parts/weight.

Subsequently, after this dispersed medium was dried and granulated, magnesium stearate was mixed by 1 part/weight. This mixture was compression-formed into tablets, each tablet having 100 mg.

(Testing on Animal)

3 beagle dogs (weight 10 - 13 kg) that were not fed were put into a group. (Testing Sample)

The following capsule was used as a testing sample.

Testing sample 1: The one prepared by filling the composition of embodiment example 1 in hard capsules (1 capsule contained 10 mg of ubidecarenone.).

Testing sample 2: Market-purchased capsule (1 capsule contained 10 mg of ubidecarenone; propylene glycol dicaprate is used for the base.)

(Method of Intake)

Each testing sample was given to the Beagle dogs (ubidecarenone 20 mg/dog) by oral intake and water 30 ml was forcibly fed to the dogs.

(Testing Method)

Blood sample collection and treatment

Blood 5 ml was collected from the front arm vein every 1, 2, 3, 4, 5, 7, and 24 hours, respectively, and the blood plasma was prepared by putting them to centrifugal separation.

(Quantification Method)

The concentration of ubidecarenone in each plasma was measured by a high speed liquid chromatography method (Ozawa et el., Arzeim Forsch vol. 36, p 689, 1986). More specifically, distilled water 0.5 ml was added to 05 ml of plasma and extracted by using 7 ml of ethanol-hexane admixture (2:5). Then, the hexane layer 4 ml was put to evaporation and dried, and dilute sulfuric acid 0.5 ml and 2% ferrous chloride 0.5 ml were added to the residue. Subsequently, this admixture was incubated at 50°C for 30 minutes, and again extracted by adding n – hexane 5ml. After acetonitrile was added to the residue, this mixture was supplied to a high speed liquid chromatigraph column. The column was 150 mm long and its diameter was 4 mm. As to the filler, TSK-Gel LS-410 (Toyo Soda was used.) was used. For the eluting solution, methanol – ethanol – acetonitrile – water (48:48:2:2) admixture was used. For detection, 273 nm UV absorption was used.

(Test Result)

Table 1 shows the average plasma concentration at a time of each blood collection from which the plasma concentration before the testing sample was given was subtracted. The tablets of the present invention demonstrated a better absorption rate for oral intake than that of reference (testing) samples.

Table 1 Plasma after each testing sample was given.

Testing sample	Concentration of ubidecarenone in the plasma: unit µg/ml									
	1 hour later	2 hours later	3 hours later	4 hours later	5 hours later	7 hours later	24 hours later			
1	0.04 (+/1 0.01)	0.33 +/- 0.02	0.30 +/- 0.02	0.43 +/- 0.07	0.44 +/- 0.09	0.44 +/- 0.08	0.22 +/- 0.09			
2	0.05 +/- 0.01	0.15 +/- 0.03	0.15 +/- 0.03	0.13 +/- 0.04	0.07 +/- 0.02	0.15 +/- 0.06	0.05 +/- 0.02			

The value in () indicates a standard deviation.

Translations
U. S. Patent and Trademark Office 3/25/03
Akiko Smith